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(54) Biaxially oriented film and magnetic recording medium comprising the same as a base film

(57) A biaxially oriented film made from a wholly aromatic polyamide produced by an acid chloride process, which contains at least one member of inorganic particles having an average particle diameter of 5 to 2,000 nm and selected from the group consisting of hydroxides, carbonates and bicarbonates of metals of groups la and III of the periodic table. The inorganic particles to derived from inorganic particles which are added as a neutralizing agent to a reaction system to neutralize hydrogen chloride by-produced by the acid chloride reaction and its content corresponds to an excess portion added to the reaction system over the amount required for neutralizing the hydrogen chloride stoichimetrically. The film is useful as a base film for a magnetic recording medium, such as a costed, a multi-layer costed, or a thin metal film high-density magnetic recording medium.

Blaxially oriented film and magnetic recording medium comprising the same as a base film

Claims of EP0913421

Translate this text

- 1. A bisasily oriented film made from a wholy sometic polyamine grounded by an acid chiorate process, which contains a feate one member of inorganic parides having an average partial disanteer of 5 to 2,000 mm and selected from the group consisting of hydroxides, carbonates and bisarbonates of metals of groups and so the product stable, the inorganic paridices being devied from inorganic paridices which are considered to the product of the product o
- 2. The biaxually oriented film of claim 1, wherein the wholly aromatic polyamide consists of at least one member of recurring units selected from the group consisting of recurring units represented by the following formulas (1) to 4) (provised that the wholly aromatic polyamide always contains (at least one of recurring units of the formulas (1) and (2)):
- wherein Ar<1> and Ar<2> are independently selected from the group consisting of
- EMI91.2
- (in which p is an integer of 0 to 4. q is an integer of 0 to 6, R is selected from the group consisting of halogen, nitro group, cyano group, sikyl group having 1 to 4 carbon atoms, alkoxyl group having 1 to 3 carbon atoms and trialityistig group) and k is 0 or a positive number).
- EMI92.1
- wherein Ar<3>. Ar<4> and Ar<5> are independently selected from the same groups as those of Ar<1> and Ar<2>, Y<1> is selected from the group consisting of -0-, -CH2-, -C(CH3)2-, -SO2-, -S- and -CO-, and i is 0 or a positive number, EMI92.2
- wherein Ar<6> is selected from the same groups as those of Ar<1> and Ar<2>, and m is 0 or a positive number, and EMI92 3
- wherein Ar<7> and Ar<8> are independently selected from the same groups as those of Ar<1> and Ar<2>.

  Y<2> is selected from the same groups as those of Y<1>, and n is 0 or a positive number.
- 3. The biaxially oriented film of claim 2 which consists of at least one member of recurring units selected from the group consisting of recurring units represented by the above formulas (1) to (4), 50 to 95 % of at least one member of aromatic groups selected from the group consisting of Ar<1> to Ar<8> having a paraorientation and forming a main chain.
- 4. The biaxially oriented film of claim 2, wherein the wholly aromatic polyamide consists of recurring units represented by the above formula (1) and recurring units represented by the above formula (2)
- 5. The biaxially oriented film of claim 4, wherein Ar<1> and Ar<2> are both a 1.4-phenylene group in the above formula (1) and Ar<3> is a 1.4-phenylene group and -Ar<4>-Y<1>-Ar<5> is
- EMI92.4 in the above formula (2)
- 6. The biaxually oriented film of claim 1, wherein the inorganic particles having an average particle diameter of 5 to 2,000 m are contained in an amount of 0.001 to 10 parts by weight based on 100 parts by weight of the wholly aromatic polyamide.
- 7. The biaxially oriented film of claim 1 which has protrusions derived from the inorganic particles having an average particle diameter of 5 to 2,000 nm, at a density of 1 x 10<1> to 1 x 10<8>/mm<2> on the surface.
- The biaxially oriented film of claim 1, wherein the wholly aromatic polyamide has a logarithmic viscosity, measured at 30 DEG C in concentrated sulfuric acid, of at least 0.5 dl/g
- The biaxially oriented film of claim 1, wherein the sum of Young's moduli in two directions intersecting at right angles on the film plane is at least 2,000 kg/mm<2>.
- 10. The biax ally oriented film of claim 1, wherein the Young's modulus in at least one direction of the two directions intersecting at right angles is at least 600 kg/mm<2>.
- 11. The biaxally oriented film of claim 1, wherein the inorganic particles have an average particle diameter of 80 to 1,000 m, protrusions derived from the inorganic particles are existent on the surface of the film at a density of 1 x 10<1> to 1 x 10<5>/mm<2>, and the sum of Young's modull in two directions intersecting at right angles on the film plane is at least 2,000 kg/mm<2>.
- 12 The biaxially oriented film of claim 11, wherein the flexural rigidity in one direction of the film plane is 3.0 to 9.5 mg.mm
- 13. The biaxially oriented film of claim 11 which has a tensile strength at break in a longitudinal direction of the film of at least 38 kg/mm<2>.
- 14. The biaxially oriented film of claim 11 which has a thermal shrinkage in one direction of the film of less than 1 %
- 15. The baxially oriented film of claim 1, wherein the inorganic particles have an average particle diameter of 5 to 150 nm, protrusions derived from the inorganic particles are existent on the surface of the film at a density of 5 x 10-3> to 1 x 10-82-/mm22-, and the sum of Young's moduli in two directions intersecting at night angles on the film plane is at least 2,000 kg/mm22>
- 16. The biaxially oriented film of claim 15, wherein elemental halogen ions are not detected from the surface of the film by photoelectric spectrophotometry.

- 17. The biaxially oriented film of claim 1 which has a thickness of 0.1 to 20 mu m.
- 18. A base film for a magnetic recording medium, which has a coating layer on at least one surface sized in the biaskally oriented film of claim 11. In coating layer confirming a water-departable complexate residence of the biaskally oriented film of claim 11. In coating layer comprising a water-departable comprised residence of the coating layer coating and the coating layer coating and a strength of the coating layer coating and a strength of the coating layer layer coating layer coating layer layer coating layer layer coating layer layer coating layer layer layer coating layer l
- 19. The base film for a magnetic recording medium of claim 18, wherein the coating layer contains fine particles having an average particle diameter of 10 to 50 nm in an amount of 1 to 50 wt% based on the above border component and satsfies the following equation: 0.2 < □ vt < □ 2.0</p>
- t is a thickness of the coating layer (nm) and
- d is an average particle diameter of the fine particles (nm).
- 20. The biaxially oriented film of claim 15 which is a base film for a magnetic recording medium.
- 21. A coated high-density magnetic recording medium which comprises the biaxially oriented film of claim 11 and a magnetic coating layer formed on one surface side of the biaxially oriented film.
- 22. A coated high-density magnetic recording medium which comprises the base film for a magnetic recording medium of claim 18 and a magnetic coating layer formed on the coating layer of the base film.
- 23. A thin metal film high-density magnetic recording medium which comprises the biaxially oriented film of claim 15 and a deposited metal thin film formed on one surface side of the film.
- 24. A multi-layer coated high-density recording medium which comprises the baxally oriented film of claim 15 and magnetic coating layers formed on one surface side of the film, the magnetic coating layers consisting of a non-magnetic buffer coating layer formed directly on the biaxally oriented film and a magnetic coating layer formed on the buffer layer.

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Biaxially oriented film and magnetic recording medium comprising the same as a base film

Description of EP0913421

Translate this text

Detailed Description of the Invention

[0001] The present invention relates to a biaxially oriented film made from a wholly aromatic polyamide and to a magnetic recording medium comprising the same as a base film. More specifically, it relates to a biaxially oriented film made from a wholly aromatic polyamide, which is excellent in handling properties such as strength, slipperiness and winding property and has excellent abrasion resistance, and to a magnetic recording medium comprising the same as a base film.

[0002] in recent years, remarkable progress has been made in high-density magnetic recording, as exemplified by the development and practical implementation of a thin layer coated magnetic recording medium in which a needle-like magnetic powder material such as metal powders or iron oxide powders is coated on a coated magnetic recording medium to a thickness of 2 pm or less. For example, an extremely thin layer coated magnetic recording medium for high-density magnetic recording is known (refer to technical report MR 94-78 issued by the institute of Electronics, Information and Communication Engineers (1995-021)

[0003] Since a coated magnetic recording medium of the prior art (a magnetic recording medium in which a mixture of magnetic powders and an organic polymer binder is coated on a non-magnetic base film) has a low recording density and a long recording wavelength, it has a large thickness of about 2 mu m or more. On the other hand, in the case of an extremely thin layer coated magnetic recording medium, there is proposed a magnetic layer as extremely thin as 0.13 mu m, though it has a non-magnetic under-layer

[0004] Along with a reduction in the size of a recording medium itself, it is desired to reduce the thickness of a base film itself in order to increase recording capacity.

[0005] An aromatic polyamide film is a material which is excellent in strength, heat resistance and the like due to high Young's moduli as a thin film and suitable for use in a high-density magnetic recording medium.

[0006] JP-A 51-129201 (the term "JP-A" as used herein means an "unexamined published Japanese patent application") discloses a magnetic recording medium which comprises a magnetic layer and an aroma polyamide film in which 70 % or more of all the bonds are para bonds and at least 15 % of phenyl nuclei have at least one substituent of halogen and nitro groups at the ortho-positions of an amide bond

[0007] In the high-density magnetic recording medium, the surface condition of a non-magnetic base film greatly influences the surface properties of a magnetic layer, and surface irregularities cause noise in recording and reproduction signals. Therefore, it is desirable that the surface of the non-magnetic base film he as smooth as nossible

[0008] On the other hand, from a viewpoint of the formation of a non-magnetic base film and handling properties such as transportation, damaging, wind-up and wind-off in the process of film formation, when the surface of the film is too smooth, slipperiness between films deteriorates, a blocking phenomenon occurs, roll formation becomes difficult, and the yield of products lowers, thereby inducing an increase in production costs. Therefore, from a viewpoint of production costs, it is desirable that the surface of the nonmagnetic base film be as rough as possible.

[0009] As means for satisfying the above antinomic requirements, there is proposed a method for forming protrusions which are appropriate in terms of height and density and derived from particles having optimal particle diameters on the surface of the base film

[0010] As a method for forming protrusions on the surface of an aromatic polyamide film, there have bee proposed a) one in which a predetermined amount of inorganic particles are added and b) one in which organic polymer particles or inorganic particles whose surface is treated with an organic polymer are added.

[0011] JP-A 61-246919 discloses a film which is essentially composed of an aromatic polyamide or aromatic polyimide containing 0.01 to 10 wt% of inorganic particles as a film produced by a method a). The film has protrusions derived from the inorganic particles and specified as follows. The heights Rm of fine protrusions range from 30 to 500 ANGSTROM, the area average diameter DA of the fine protrusions ranges from 0.01 mum or more to less than 0.3 mum, the average flatness DA/Rm of the protrusions is 15 or less, the number NA of fine protrusions having a maximum diameter of 0.01 mu m or more is 10<4> to 10<9>/mm<2>, the number of fine protrusions having a long diameter/short diameter ratio a/b (a: maximum glameter of fine protrusion, b: diameter intersecting the maximum diameter at a right angle) of 2 or more is 0.2 x NA or less, and the number of fine protrusions having a circularity pi DN/CN (DN: area diameter of fine protrusion. CN: circumference of fine protrusion, pi : ratio of the circumference of a circle to its diameter) of 0.7 or less is 0.2 x NA or less

[0012] JP-A 8-203064 discloses a film made from an aromatic polyamide or an aromatic polyimide, which contains organic polymer particles having an average particle diameter of 10 to 500 nm or inorganic particles surface-treated with an organic polymer, has a center line surface roughness Ra of one surface of 0.5 to 50 nm and an Ra of the other surface of 10 nm or less

[0013] However, the above methods using externally added particles involve such a defect that the particles fall off from the protrusion on the film surface by friction between the guide pin and the surface of the film in the calendering step after the coating of a magnetic layer or during actual use and contaminate the step because the externally added particles easily applomerate in the added slurry

[0014] Particularly, in the calendering step after the application of a magnetic coating, extremely high shearing force is applied to the base film. Therefore, large voids formed around the agglomeration of the particles produce a great amount of abrasion dust due to the fall-off of the particles, thereby contaminating the calender roll with the result of a great reduction in productivity

[0015] It is an object of the present invention to provide a novel biaxially oriented wholly aromatic polyamide

[0016] It is another object of the present invention to provide a blaxially oriented wholly aromatic polyamide film for a magnetic recording medium, which is excellent in handling properties such as strength slipperiness and winding property and has high abrasion resistance

[0017] It is still another object of the present invention to provide a magnetic recording medium which comprises the biaxially oriented wholly aromatic polyamide film of the present invention as a base film, has excellent electromagnetic conversion charactanstics and is suitable for high-dansity recording.

[0018] Other objects and advantages of the present invention will become apparent from the following description

[0019] According to the present invention, firstly, the above objects and advantages of the present invention can be attained by a biaxially oriented film made from a wholly aromatic polyamide produced by an acid chloride process, which contains at least one member of inorganic particles having an average particle diameter of 5 to 2,000 nm and selected from the group consisting of hydroxides, carbonates and bicarbonates of metals of groups is and ills of the periodic table, the inorganic particles being derived from inorganic particles which are added as a neutralizing agent to a reaction system to neutralize hydrogen chloride by-produced by the acid chloride reaction and its content corresponding to an excess portion added to the reaction system over the amount required for neutralizing the hydrogen chloride stoichiometrically

[0020] The wholly aromatic polyamide constituting the biaxially oriented film of the present invention is not particularly limited as far as its main chain comprises an aromatic nucleus and an amide bonding group as main constituents

[0021] Preferrad examples of the recurring unit forming the wholly aromatic polyamide of the present invention include units represented by the following formulas (1) to (4):

wherein Ar<1> and Ar<2> are independently selected from the group consisting of EMIS 2

(in which p is an integer of 0 to 4, q is an integer of 0 to 6, R is selected from the group consisting of halogen, nitro group, cyano group, alkyl group having 1 to 4 carbon atoms, alkoxyl group having 1 to 3 carbon atoms and trialky/silvl group) and k is 0 or a positive number.

wherein Ar<3>, Ar<4> and Ar<5> are independently selected from the same groups as those of Ar<1> and Ar<2>, Y<1> is selected from the group consisting of -O-, -CH2-, -C(CH3)2-, -SO2-, -S- and -CO-, and 1 is 0 or a positive number,

EMI6 2 wherein Ar<6> is selected from the same groups as those of Ar<1> and Ar<2>, and m is 0 or a positive number, and

FMI6.3 wherein Ar<7> and Ar<8> are independently selected from the same groups as those of Ar<1> and Ar<2>. Y<2> is selected from the same groups as those of Y<1>, and n is 0 or a positive number.

[0022] It is preferred that the wholly aromatic polyamide in the present invention preferably contains at least one of the recurring units (1) and (2). That is, the wholly aromatic polyamide consists of (i) only the recurring units of the above formula (1), (ii) only the recurring units of the above formula (2) or (iii) only the recurring units of the above formulas (1) and (2). Further, a wholly aromatic polyamida consisting of (iv) at least one of the recurring units of the above formulas (1) and (2) and at least one of the recurring units of the above formulas (3) and (4) is preferred

[0023] The wholly aromatic polyamide in the present invention is produced by an acid chloride process

[0024] In the abova formula (1), Ar<1> and Ar<2> are a residual group obtained by removing an acid chloride group from an aromatic dicarboxylic acid chloride and a residual group obtained by removing an amino group from an aromatic diamine, respectively.

[0025] Ar<1> and Ar<2> are independently a residual divalent aromatic group selected from the group consisting of **EMI7 1** 

[0026] In the above formulas, R is halogen, nitro group, cyano group, alkyl group having 1 to 4 carbon atoms, alkoxyl group having 1 to 3 carbon atoms or trialkylsilyl group, p is an integer of 0 to 4 and q is an integer of 0 to 6

[0027] Illustrative examples of the abova residual divalent aromatic group include 1,4-phanylene group, 1,3phenylene group, 1,2-phenylene group, 4,4'-biphenylene group, 4,3'-biphenylene group, 4,2'-biphenylene group, 3,3'-biphenylene group, 3,2'-biphenylene group, 1,4-naphthylene group, 1,5naphthylene group, 1,6-naphthylene group, 2,6-naphthylene group, 2,7-naphthylena group and groups obtained by replacing these groups by the above substituants. Of these, 1,4-phenylene group, 4,4biphenylene group, 1.4-naphthylene group, 2.6-naphthylene group and groups obtained by replacing these groups by the above substituents are preferred

[0028] Preferred examples of the halogen represented by the substituent R include fluorine, chlorine and bromine. The alkyl group having 1 to 4 carbon atoms may be straight-chain or branched-chain and preferred examples thereof include methyl group, ethyl group, n-propyl group, isopropyl group, n-butyl group, isobutyl group, t-butyl group and the like. The alkoxyl group having 1 to 3 carbon atoms may be straight-chain or branched-chain, as exemplified by methoxy group, ethoxy group, n-propoxy group and isopropoxy group. Further, the trialkylsilyl group is preferably a trialkylsilyl group each alkyl group of which independently has 1 to 3 carbon atoms, such as trimethylsilyl group.

[0029] Illustrative examples of the recurring unit represented by the above formula (1) are obvious from the

above specific examples of Ar<1> and Ar<2>. For example, when Ar<1> and Ar<2> are both a 1,4phenylene group, the recurring unit is represented by the following formula:
EMIS 1

wherein k is the same as defined in the above formula (1).

[0030] In the above formula (2), Ar<3>, Ar<4> and Ar<5> are independently selected from the same groups as those of Ar<1> and Ar<2> Y<1> is -O-, -CH2-, -C(CH3)2-, -SO2-, -S- or -CO-.

[0031] Illustrative examples of the recurring unit represented by the above formula (2) are obvious from specific examples of Arc19, Arc29 and Y<19 for example, when Arc39, Arc49 and Arc59 are all a 1,4-EMI8 2.

EMI8 2.

wherein I is the same as defined in the above formula (2)

[0032] In the above formula (3), Ar<6> is selected from the same groups as those of Ar<1> and Ar<2>

[0033] Illustrative examples of the recurring unit represented by the above formula (3) are obvious from specific examples of Ar<5 = And Ar<2k. For example, whan Ar<6b is a 1.4-phenylene group, the recurring unit is represented by the following formula:

wherein m is the same as defined in the above formula (3)

[0034] In the above formula (4), Ar<7> and Ar<8> are independently selected from the same groups as those of Ar<1> and Ar<2>, and Y<2> is selected from the same groups as those of Y<1>.

[0035] Illustrative examples of the recurring unit represented by the above formula (4) are obvious from specific examples of Ar-19, Ar-22 and Y-15 For example, when Ar-72 and Ar-18-3 are both a 1,4phenylene group and Y<2> is -O., the recurring unit is represented by the following formula. EMIS 2

wherein n is the same as defined in the above formula (4)

[0039] The wholly aromatic polyamide in the present invention preferably consists of at least one member of recurring units selected from the group consisting or recurring units represented by the above formulas (1) to (4), and 50 to 99.5 % of at least one member of aromatic groups selected from the group consisting of ARX-10 ARX-9 selected in at least one member of the selected recurring units has a para-orientation and forming a main chain. In this case, the wholly aromatic polyamide contains at least one of the recurring units recreasemed by the above formulas (1) and (2).

[0037] In the present invention, the aromatic group having para-orientation refers to, for example, 1,4-phenylane group, 4,4'-biphenylane group, 1,4-naphthylene group or 2,6-naphthylene group.

[0.038] From the viewpoint of the recurring units, the wholly aromatic polyamide in the present invention preferably consists of recurring units represented by the above formulae (1) and occurrent of preferably recurring units represented by the above formulae (1) in which x+t> and x+t2 are both is 1.4-phenylene group and recurring units represented by the above formulae (3) in which x+t> is 4.5-phenylene group. X+t0 is -0.2-and Art<0> is 1.3-phenylene group, x+t0 is -0.2-and Art<0> is 1.3-phenylene group, and the recurring unit is regressented by the following formulae.

[0039] The blookly criented firm of the present invention comprises he above wholly anomalic polyanties as base and contains inorganic particles having an average particle disarter of 5 to 2,000 mm. The inorganic particles are selected from the group consisting of hydroxides, carbonates and bicarbonates of metals of groups I and II of the periodic table Illustrative examples of the inorganic particles include magnetism hydroxide, magnetism carbonate, carbonate, carbonate, brother carbonate, carbonate,

[0040] The inorganic particles are derived from the hydroxide, curbonate or biachtonate of a metal of group or I of the periodic table, which is added to a reaction system to neutralize hydrogen chloride by produced in an add chloride nearciton for producing the wholly aromatic polyamide, that is, a reaction between an exemptic disturbing clad delibride and an aromatic diament, in other words, by adding the inorganic particles to the reaction system in an amount exceeding the stockhometric amount of the integratic particles on the reaction system in an amount exceeding the stockhometric amount of the inorganic particles and the reaction system in an amount exceeding the stockhometric amount of the inorganic particles added to make the present invention more comprehensible, a relatedy small particle and sed of the make the present invention more comprehensible, an adaletely small particle may be comprehend consumed by a neutralization reaction but an one portion of the granticle may set the emptice of the neutralization reaction. In we do not profit of the particle may set the emptice of the neutralization reaction but a make the inorpanic particles and the demandration of the neutralization reaction. In we do not profit of the particle may set the emptice of the neutralization reaction but a make the inorpanic particles as when a new setting the particle dispatch is not provided to the particle may set the emption of the neutralization of the neutralization in excellent in the film, the reaction system is preferrably in the reaction. If you do not make the inorpanic particle and the setting of the neutralization reaction but as the make the inorpanic particle and the setting of the neutralization reaction.

[0041] As described above, the inorganic particles contained in the biasisty oriented film of the present intention are obtained as the residual particles of the neutrations agent that hey ophiretization step of the wholly aromatic polyamide. Since they are fully dispersed in the system in the polymetization test, there is a particle allower for the present particle allower for the particles contained in the film is installed than 5 million protrusions on the surface its small and excellent handling properties such as slipperiness and winding protrusions on the surface its small and excellent handling properties under such as simperiness and winding protrusions on the surface its small and excellent handling properties under a surface of the present particle statement is larger than 2.000 nm. volds formed by structuring the film become too larger, thereby deteriorating abrassion reasstance professional processing and the properties of the properties of the present particle statement of the present particles statement in larger than 2.000 nm. properties and processing statement and the processing preferrably 800 not 000 nm. processing preferrably 800 not 000 nm.

[0042] The average particle diameter of the inorganic particles to be added to the reaction system is preferably 150 to 20,000 nm, more preferably 150 to 15,000 nm, as described above. When the average particle diameter is smaller than 150 nm, their effect of forming protrusions on the surface is small and excellent handling properties such as slippenness and winding property cannot be obtained

- disadvantageously. When the average particle diameter is larger than 20,000 nm, the particle diameters of the residual particles after the neutralization reaction are large, and voids formed by stretching the film become too large, thereby deteriorating abrasion resistance disadvantageously.
- [0043] Means of controlling the particle diameters of the inorganic particles is preferably a method in which the particles are dispersed in a solvent and then, ground using such an apparatus as a sand grinder and the obtained dispersion is filtered to remove large particles.
- [0044] The wholly aromatic polyamide in the present invention is produced by an acid chloride process. Polymerization may be carried out by an interfacial polymerization or solution polymerization. Of these, solution polymerization is preferred.
- [0045] Preferred polymerization solvents which can be used in the solution polymerization include dimethylformamide, dimethylacetamide, N-methylpyrrolidone, N-methylcaprolactam, dimethyl sulfoxide, hexamethylphosphoryl triamide, letramethyl urea and 1,3-dimethyl-2-imidazoidinone. They may be used alone or in combination of two or more. The polymerization solvent as a main component may be used in conjunction with other solvent.
- [0046] To the polymerization solvent may be added an inorganic salt such as calcium chloride or lithium chloride in a suitable amount before, during or after polymerization to improve the solubility of the resulting polymer.
- [0047] Acid components and amine components are reacted with each other substantially in aquimolar amounts, but either one of the components may be used excessively to control the degree of polymerization.
- [0048] A small amount of a monofunctional acid component or amine component may be used as a terminal capping agent.
- [0049] An aliphatic or aromatic amine or quaternary ammonium salt may be added to a polymerization system to capture hydrogen chloride produced by the reaction.
- [0050] Further, additives such as an ultraviolet absorber, dye, release agent and the like may be added in limits not prejudicial to the effect of the present invention.
- [0051] After the reaction is substantially completed, the above inorganic particles having an average particle of significant for 150 to 20,000 nm are added for a rection system to carry out a neutralization reaction. At this point, the inorganic particles are used in excess of an amount over the stoichiometric amount required for neutralization as described above.
- [0052] The production method of the wholly aromatic polyamide per se is detailed in JP-B 52-39719 (the term "JP-B" as used herein means an "examined Japanese patent publication").
- [0053] The description of this publication is included as the description of this specification.
- [0054] The wholly aromatic polyamide in the present invention may contain additives such as an antioxidant, antistatic agent and release agent, and other polymers in limits which do not impair the physical properties of the film.
- [0055] The wholly aromatic polyamide in the present invention preferably contains inorganic particles having an average particle diameter of 5 to 2,000 nm in a mount of 0.001 to 10 parts by weight, more praferably 0.004 to 8 parts by weight, particularly preferably 0.01 to 5 parts by weight, based on 100 parts by weight of the wholly aromatic polyamide.
- (0056) The wholly aromatic polyamide in the present invention perferably has a logarithmic viscosity, measured at 30 GEC on concentrated sulfuric acid, or let pest 0.5 d/ig, more perfeatably at least 1.0 d/ig. The logarithmic viscosity of the polymer before film formation is obtained by measuring at 30 DEC C in concentrated sulfuric acid the polymer which has been emp-precipitated and separated after a polymer stock solution containing the wholly aromatic polyamide after polymerization has been added to a solvent such as an alcohol or vasting.
- [0057] The biaxially oriented film of the present invention can be produced as follows
- [0.055] After polymerization is carried out as described above, the polymer stock solution containing the wholly aromate polymeric can be added to a solvent such as an alcohol of variety, and the polymer can be re-precipitated, separated and then dissolved in a solvent again so that it can be used for the formation of fifth. However, the polymer stock solution is preferably used directly or upon algorithment to an appropriate first However, the polymer stock solution is preferably used directly or upon algorithment to an appropriate polymerization solution with a solvent. As the solvent may be used the same as those fitsed for the polymerization solvent.
- [0059] The polymer stock solution prepared as described above is formed into a film by a solution film formation process. The solution film formation process is selected from dry-wet, dry and wet processes. Dry-wet and topy processes are preferred because a film having excellent surface properties is obtained.
- [0050] When a film is formed by a wet process, the stock solution is preferably extruded into a film formation that directly from a nozicy or extruded onto a support such as a dum and hen introduced into a wet process bath together with the support. This bath is generally an aqueous medium and may contain an organic solvent or morganic sate by seasing through the wet process bath, the contained safe or organic solvent can be extracted. The time required to pass through the wet process bath is preferably 10 seconds to 30 minutes. Brown bit differs according to the thickness of a film.
- [0061] The polymer passed through the wet process bath is stretched in a longitudinal direction, dried, stretched in a transverse direction and heat set. These treatments are preferably carried out at 100 to 500 DEG C for a total time of 1 second to 30 minutes.
- [0062] When a film is formed by a dry-wet process, the stock solution is extruded onto a support such as a drum or endless belt to form into a thin film, a solvent is scattered from the thin film layer, and the film is

dried until it achieves self-holding properties. Preferred drying conditions are a temperature of room temperature to 300 DEG C and a time of 60 minutes or less. The film after the during step is separated from the support, introduced into a wel step to remove the salt and solvent from the film, stretched, dried and heat self, like the wet process.

[0063] When the dry process is employed, the film which has been dried on a drum or endless belt and achieved self-holding properties is separated from the support, dried to tremove the residual solvent, stretched and heat set. These treatments are preferably carried out at a temperature of 100 to 500 DEG C for a total time of 1 second to 30 minutes

(0064) The film is preferably stretched at an area stretch ratio of 5.0 times or more, more preferably 6.0 to 10.0 times.

[0065] The thickness of the biaxially oriented film of the present invention obtained by the above processes is not particularly limited but preferably 0.1 to 20 mu m. more preferably 0.5 to 10 mu m.

[0066] When the thickness of the film is within the above range, it shows excellent handling properties such as strength, slipperiness and winding property and high abrasion resistance, and can be advantageously used for a magnetic recording medium.

[0607] The bisasily oriented film of the present invention preferably his profusions derived from the inorganic particles having an average particle dameter of 5 to 2,00 mm, at a density of 1 x 100-15 to 1 x 104-80-imm22+, more preferably 1 x 104-15 to 1 x 104-80-imm22+, on the surface. When the density of this profusion is lower than 1 x 104-15-imm22+, fristion coefficient becomes large, whereby good running properties are thangly obtained, winding property readly deterioristics and blocking between films is liable to selectionage. The conversion or particle that is readly deterior and et also sharp the conversion or particle that is readly deterior and et also sharp the conversion or particle that is readly deterior and et also sharp the conversion.

[008] The sum of Young's moduli in two directions intersecting at right angles (for example, a longitudinal direction and a stansverse direction) on the plane of the bissally oriented film of the present invention is praferably at least 2,000 kg/mm<sup>2</sup>. This preferred upper limit of the sum of Young's moduli is 4,000 kg/mm<sup>2</sup>. This preferred upper limit of the sum of Young's moduli is 4,000 kg/mm<sup>2</sup>. When the sum of Young's moduli is 4,000 kg/mm<sup>2</sup>. When the sum of Young's moduli is 4,000 kg/mm<sup>2</sup>. When the sum of Young's moduli is also than 2,000 kg/mm<sup>2</sup>. And this film is used as a base film for a magnetic recording medium, the contact between the magnetic recording medium and amagnetic head becomes radequate, whenly electromagnetic conversion characteristics are liable to deteriorate and a base film having a thickness in the order of several micrometers cannot have sufficient streety disadvardingsome.

[0069] The Young's modulus in at least one of the two directions intersecting at right angles is preferably at least 600 kg/mm<2>.

[0070] The following two embodiments of the biaxially oriented films of the present invention are particularly

(i) A biaxally criented film containing incrganic particles having an average particle diameter of 80 to 1,000 m and having profrusions derived from the inorganic particles at a density of 1 x 10<15 to 1 x 10<15 /mm<2> on the surface and a sum of Young's moduli in two directions intersecting at right angles on the film plane of at least 2,000 kg/mm<2>.

(ii) A biaxally oriented film containing inorganic particles having an average particle diameter of 5 to 150 nm and having protrusions derived from the inorganic particles at a density of \$ 1.045 to 1 x 10-85 mm <2> on the surface and a sum of Young's moduli in two directions intersecting at right angles on the film plane of at least 2,000 kg/mm <2>

(0071) The basisity oriented film (i) preferably has a flexural rigidity of 3.0 to 9.5 mg mm, more preferably 50 to 9.5 mg mm, in one direction on the film plane. When the flexural rigidity is one direction, for example in a longitudinal direction is less than 3.0 mg mm or more than 9.5 mg, mm, the contact state between the film and the recording head is act to be inadequate, thereby deteriorating electromagnetic conversion characteristics disadvantageously.

[0072] Further, the tensile strength at break of the film in a longitudinal direction is preferably 38 kg/mm<2> or more, more preferably 42 kg/mm<2> or more. When the tensile strength at break is less than 38 kg/mm<2> film can hardly be used as a tape which runs and stops repeatedly.

[0073] The thermal shrinkage at 150 DEG C in one direction of the film is preferably less than 1 %. When the thermal shrinkage is 1 % or more, the reliability of the performance of the film is liable to be lost depending on the use environment of a recording medium disadvantageously.

[0074] Further, in the above biaxially onented film (iii), elemental halogen ions are not detacted from the surface of the film by photoelectric spectrop-totmetry. This biaxially oriented film is free from the damage of the magnetic metal thin film by elemental halogen even in a high-temperature and high-humdidy environment and can retain its performance when it is used in a magnetic recording medium of a metal thin film tune.

[0075] The biaxially oriented film of the present invention can be used by itself as a base film for a magnetic recording madium per se and it is also used as a base film for a magnetic recording medium after a coating layer is formed on the film.

[0076] It is particularly advantageous that the biaxially oriented film (i) be used as a base film after a coating layer is formed thereon.

[0072] Therefore, according to the present invention, there is also provided a base film for a magnetic recording medium, which has on a flactor to esurface of the hazardy oriented firm (i) of the present invention or a coating layer comprising as a binder component a valler-diagnesible copolyster reain which comprises 40 or 98 molfs of 26-perhythereinderican-topic acid, 0 10 to find's of an anomatic distancy calc dampting a sulfonate salt group and 0 to 55 molfs of other aromatic disastrocyte acid as the anomatic disastrocyte acid components and 40 to 10 molfs of 40-perhips gloyed 10 to 81 molfs of 40 molfs of 40 molfs or 40 molfs or

#### W/R r of 5 to 200 nm

[0078] in the composition of the water-dispersible copolyester resin as the binder component of the coating layer, when the proportion of 2,6-naphthalenedicarboxylic acid is smallar than 40 mol%, the blocking resistance of the film lowers disadvantageously, while when the proportion is larger than 90 mol%, the copolyester resin is difficult to dissolve in a hydrophilic organic solvent in the step of dispersing the copolyester resin in water, theraby making difficult water dispersion. In this case, it is effective and preferred to improve non-crystallinity by copolymerizing glycol components. However, when the proportion of 2,6naphthalenadicarboxylic acid is larger than 99 mol%, even if glycol components for improving noncrystallinity are copolymerized, the copolyester resin does not dissolve in a hydrophilic organic solvent any longer. Further, when the proportion of the aromatic dicarboxylic acid having a sulfonate salt group is smaller than 0.1 mol%, the hydrophilic nature of the copolyester resin deteriorates, thereby making difficult water dispersion disadvantageously. When the proportion is larger than 5 mol%, the blocking resistance of the film deteriorates disadvantageously

[0079] Preferred examples of the aromatic dicarboxylic acid having a sulfonate salt group include 5-sodium sulfoisophthalic acid, 5-potassium sulfoisophthalic acid, 5-lithium sulfoisophthalic acid and 5-phosphonium sulfoisophthalic acid. For improving water dispersibility, alkaline metal salts such as 5-sodium sulfoisophthalic acid, 5-potassium sulfoisophthalic acid and 5-lithium sulfoisophthalic acid are more preferred

[0080] The acid components of the copolyester resin include 2,6-naphthalenedicarboxylic acid and an aromatic dicarboxylic acid having a sulfonate salt group in the abova proportions, and other aromatic dicarboxylic acid may be used in combination with these. Illustrative examples of the other aromatic dicarboxylic acid include terephthalic acid, isophthalic acid, phthalic acid, biphenyldicarboxylic acid and the like. Of these, isophthalic acid is particularly preferred.

[0081] When the proportion of ethylene glycol as a glycol component of the copolyester resin is smaller than 40 mol%, the blocking resistance of the film lowers disadvantageously.

[0082] The addition product of bisphenol A with a lower alkylene oxide which can be used in conjunction with ethylene glycol is a compound represented by the following formula: FMI211

wherein X is H or lower alkyl having 1 to 5 carbon atoms, and I + m is an integer of 2 to 10. The use of the addition product of bisphenol A with a lower alkylene oxide is effective in improving the water dispersibility of the copolyester. Illustrative examples of the lower alkyl represented by X in the above formula include alkyl groups having 1 to 5 carbon atoms such as methyl, ethyl, propyl, butyl and pentyl. Of these, methyl is particularly preferred. The compound is particularly preferably an addition product of bisphenol A with propylene oxide. When the value of I + m increases, the blocking resistance of the copolymer lowers. Therefore, I + m is preferably as small as possible and the most preferably 2.

[0083] The additional product of bisphenol A with a lower alkylene oxide is preferably used in an amount of 10 to 50 mol%, more preferably 20 to 40 mol%, based on the total of all glycol components.

[0084] The glycol components of the copolyester include ethylene glycol and an addition product of bisphenol A with lower sikylene oxide in the above proportions. When the total proportion of these glycol components is smaller than 10 mol%, other aliphatic or alicyclic glycol can be used in combination with these. Preferred examples of the other alignatic or alloyellic glycol include 1.4-butanediol. 1.4cyclohexanedimethanol and the like

[0085] The coating layer may contain fine particles having an average particle diameter of 10 to 50 nm in addition to the above binder component. The average particle diameter of the fine particles is preferably 15 to 40 nm. When the average particle diameter is smaller than 10 nm, the blocking resistance becomes insufficient, while when the average particle diameter is larger than 50 nm, the coating layer must be made thick to prevent the particles from falling off with the result that blocking resistance becomes insufficient.

The content of the fine particles is 1 to 50 wt%, preferably 1 to 35 wt%, more preferably 1 to 25 wt% based on the weight of the copolyester resin

[0086] Preferably, the fine particles have a relatively low specific gravity so that they hardly sediment in a coating solution. Preferred examples of the fine particles include fine particles of heat resistant polymers (such as cross-linked silicone resin, cross-linked acryl resin, cross-linked polystyrene, melamine formaldehyde resin, aromatic polyamide resin, polyamidelmide resin, cross-linked polyester, wholly aromatic polyester and the like), silicon dioxide (silica) and calcium carbonate. Of these, cross-linked silicone resin fine particles, silica and core-shell typa organic fina particles (core: cross-linked polystyrena, shell: polymethyl methacrylate) are particularly preferred

[0087] When the above fine particles are contained, the thickness of the coating layer desirably satisfies the following equation (5): "/5\" n 2 </= t/d </= 2 n

wherein t is a thickness (nm) of the coating layer, and d is an average particle diameter (nm) of the fine particles

preferably the following equation (6):

"(6)" 0.3 </= t/d </= 1.5 wherein t and d are the same as defined in the equation (5).

more preferably the following equation (7): "(7)" 0 4 </= t/d </= 1.0

wherein t and d are the same as defined in the equation (5).

particularly preferably the following equation (8): "(8)" 0.5 </= t/d </= 0.8

wherein t and d are the same as defined in the equation (5).

[0088] When t/d (thickness of coating layer/average particle diameter of fine particles) is less than 0.2 the fine particles cannot be held by the binder and fall off in the step of forming a film or producing a magnetic tape so that many drop outs are produced and electromagnetic conversion characteristics are deteriorated. When t/d is more than 2.0, the number of particles projecting from the binder is reduced and sufficient blocking resistance cannot be obtained

(0089) it is preferred that the fine particles should not applomerate in the coation layer, and it is further

preferred from the viewpoint of abrasion resistance that the density of agglomerates of 10 or more primary particles is 0 to 30/10 mu m<>.

(0090) The center plane average roughness WRs of the surface (not in contact with the biaxially onented film) of the coatility payers 0.5 to 10 m, preferably 1 to 1 m, more preferably 1.5 to 5 m. When MRs is less than 0.5 mm, the slippenness between the film and the pass roll degrades in the step of forming a film producing a magnetic tape, the film is chipped so that many drop outs are produced and electromagnetic conversion characteristics deteriorate. When WRs is more than 10 mm, the magnetic surface becomes rough and self-story electromagnetic conversion characteristics can only the obligated.

[0091] The 10-point average roughness WRz of the surface of the above film is 5 to 200 mm, preferably 10 to 150 mm, more preferably 15 to 100 nm. When WRz is less than 5 nm, the degassing property of the film to be wound on a roll degrades to make it difficult to obtain a good roll shape of the film, thereby greatly reducing product yeld. When WRz is more than 200 nm, the magnetic surface becomes rough and satisfactory electromagnetic conversion characteristics cannot be obtained.

[0092] Coating for the formation of the coating layer can be carried out during or after film formation but preferably during film formation. Coating is preferably carried out after stretching and heat setting to prevent the excessive heating of the coating layer.

[0093] Coating may be carried out in accordance with a commonly used method, such as kiss coating, reverse coating, gravure coating, die coating or the like;

[0094] The thickness of the coating layer is 1 to 50 nm, preferably 5 to 35 nm. When the thickness is smaller than 1 nm, adhesion becomes insufficient, while when the thickness is larger than 50 nm, blocking resistance lowers. When the coating layer contains fine particles, it is desired that the relationship between the average particle dismeter of the fine particles and the thickness of the coating layer should satisfy the above equations (5) to (8) as described above.

[0095] It is possible to contain other components in the coating layer as required in limits not prejudicial to the effect of the present invention for the purpose of improving slipperiness and preventing electrification.

[0096] A base film for a magnetic recording medium, which comprises the above biaxially oriented film (i) provided by the present invention and a coating layer on at least one surface side of the film (i) provides a coated high-density magnetic recording medium by forming a magnetic coating layer on the coating layer.

[0097] That is, according to the present invention, there are also provided a coated high-density magnetic recording medium which comprises the above biaxially oriented film (i) of the present invention and a magnetic coating layer formed on one side of the film (i) and a coated high-density magnetic recording medium which comprises the above base film for a magnetic recording medium having a coating layer on the blaxially oriented film (ii) and a magnetic coating layer formed on the coating layer.

[0098] Magnetic coatings include metal-based magnetic coatings and metal oxide-based magnetic coatings.

[0099] The metal-based magnetic coatings are prepared by containing a magnetic metal or a magnetic material essentially composed of a magnetic metal in a binder, as exemplified by a coating prepared by uniformly dispersing from or needle-like magnetic fine powders essentially composed of iron in vinyl chloride or vnyl chloride.

[0100] The metal oxide-based magnetic coating is prepared by containing a magnetic metal oxide or a magnetic material essentially composed of a magnetic metal oxide in a binder, as exemplified by a coating prepared by uniformly dispersing needle-like magnetic fine powders such as iron oxide or chromium oxide, or leaf-like magnetic fine powders such as barium ferrite in a binder such as vinyl chloride or vinyl chloridevinyl acotate copylmer

[0101] The magnetic coating is applied to ensure that the thickness of the obtained magnetic layer should be 2 mu m or less, preferably 0.1 to 2 mu m.

[0102] A coated magnetic recording medium for high-density recording which has excellent electromagnetic conversion characteristics such as output at a short-wavelength range, SN and CN, few drop outs and a small error rate can be obtained by forming a back coat layer on a side opposite to the magnetic layer in accordance with a known methy and converse the converse of the conver

[0103] Similarly, the above blaxially oriented film (ii) of the present invention can provide a high-density magnetic recording medium.

(0.04) That is, according to the present invention, there are also provided a metal thin film high-density magnetic recording medium which comprises the above bisaclay oriented filling (o) of the present invention and a deposted metal thin film on one surface size of the film (0), and a multi-layer coated high-density magnetic recording medium which compress the above bisaclay covered filling (o) of the present invention compressed that the present invention of the present invention consistent of a non-magnetic buffer coating layer formed on this surface of the biaxisity one next fill mixetly and a magnetic corting layer formed on the buffer coating layer.

[0.05] The metalt bin fifth high-density magnetic recording medium is produced by forming a ferroelectric metalt bin fifth liquid made from inco clouds, chromium or an alloy or node thereto by avouum deposition, sputieting, lon plating or the like on one surface of the blackally oriented fifth (0), a producive layer of diamond-like carbon (0). C) or the like and a fluorine-contribung carbonyile active based lubricant layer on the surface of the ferroelectric metal thin fifth layer in the mentioned order according to purpose and application and as received and untime a fluorine because the coolsist both the metalt high fifth layer.

[0106] This metal thin film high-density magnetic recording medium which has excellent electromagnetic conversion characteristics such as output at a short-wavelength range, SN nand CN, few drop outs and a small error rate is extremely useful as a tape medium for Hi8 for analog signal recording, digital video cassette recorders (DVC) for digital sooial recording, data 8 mm and DSIV.

[0107] The multi-layer coated high-density magnetic recording medium is made multi-layered by dispersing

- titanium oxide fine particles in the same organic binder as that for the magnetic layers and applying the organic binder as a non-magnetic layer and the magnetic coating to at least one side of the biaxially oriented film (ii) of the present invention in this order.
- [0108] The magnetic coating includes metal-based magnetic coating and metal oxide-based magnetic coating. The metal-based magnetic coating and the metal oxide-based magnetic coating are the same as those listed above.
- [0109] The thickness of the magnetic leyer is preferably 1.1 mu m or less, more preferably 0.1 to 1.0 mu m.
- [0110] The thickness of the non-magnetic layer is preferably 0.1 to 2.0 mu m, more preferably 0.2 to 1.5 mu m.
- [0111] A costed magnetic recording medium for high-density recording which has excellent electromagnetic conversion characteristics such as output at a short-wavelength range. SN and CN, few drop outs and a small error rate can be obtained by forming a back coat layer on a side opposite to the magnetic layer in accordance with a known method.
- [0112] This multi-layer coated high-density magnetic recording medium is extremely useful as a tape medium for 8 mm viseo, Hilb. Seat-cam SP and MV-HS for analog signal recording, digital video cassette recorder (DVC) for digital signal recording, data 8 mm, DDSIV. digital beta -cam, D2, D3, SX and data streamer DIC. for digital signal recording.
- [0113] The above W-VHS is a VTR for analog HDTV signal recording, and the DVC can be used for digital HDTV signal recording. It can be said that the aromatic polyamide film of the present invention is a base film extremely useful for a magnetic recording medium for VTRs for HDTV signal recording.
- [0114] The following tox samples are given to further illustrate the present invention, it is to be understood that he present invention is not limited by the following examples and various changes and modifications may be made in the present invention without departing from the spirit and scope thereof. Physical properties and characteristic properties in the present invention were measured and defined as follows.
- (1) pH of polymer solution
- [0115] After 5 g of a polymer solution was collected, edded to 300 ml of water and stirred for 2 minutes, pH of the resulting aqueous solution was measured.
- (2) logarithmic viscosity
- [0116] A polymer stock solution containing an aromatic polyamide after polymerization was added to a solvent such as an alcohol or weter, and the logarithmic viscosity of a polymer which was re-precipitated and separated was measured at 30 DEG C in concentrated sulfuric acid.
- (3) area stretch ratio
- [0117] This was obtained by multiplying a stretch ratio in a longitudinal direction by a stretch ratio in a transverse direction of the film
- (4) average perticle diameter of particles
- (a) average particle diameter of particles before addition to polymer solution
- [0118] Particles were measured for diameters using the centrifugal particle size analyzer Model CP-50 of Shimadzu Corporation. A particle diameter equivalent to 50 weight percent was read from a cumulative curve of diameters of the indivioual particles and the amount thereof celculated based on the obtained centrifugal sedimentation curve, and the obtained value was taken as an average particle diameter (refer to "Perticle Size Measurement Technique" issued by Nikara Koypo Press. p. 24:2247, 1930.
- (b) average perticle diemeter of perticles contained in film
- [0119] The film was dissolved in N-methylpyrrolidone (may be abbrevieted as NMP hereinafter) containing 3 with of lithium chloride by stirring at 100 DEG C for 30 minutes and the residual film undissolved in NMP was measured in the same manner as in (a).
- [0102] For confirmation, a similar sample prepared by encapsulating a small piece of the film in an epoxy rean (Epomount of Refinetes Co. Lud., and the silicon source of the film in support with the resi using the Microtore 255 of Reicher-Lung Co. Lud., and the silicon was observed by a transmission electron microscope (H-800 of Histan, Lud.) to obtain the particle insimetter of 25 cross sections of the particles A. The obtained particle dismetters were averaged to obtain an average particle diameter. As a result, the same value as a shown was obtained.
- (5) density of protrusions
- [0121] 25 Photos of the surface of the film were taken at random at an angle of 45 DEG and a magnification

of 5,000X using a scanning electron microscope (T-300 of JEOL Ltd.) to count the number of protrusions on the surface of the film. The number of protrusions per 1 mm<2> was calculated from the average of the count values.

(6) void ratio

[0122] The film was eiched to a depth of 500 mm from the surface using a pisama reactor [PR3 of Varianot Agalau Co., Liu, 32 Photos of the surface of the ethods angine verte taken it random at an angle of 0 DEG and a magnification of 5,000X using a scanning electron microscope (T-300 of JEC). Ltd.) to measure the long diameter of a particle which appeared on the surface and the long diameter of a void around the particle. The void ratio was obtained from the following equation:

(7) mechanical properties

(a) Young's moduli

[0123] A sample was prepared by cutling the fifth to a width of 10 mm and a length of 150 mm, and this sample was pulled by a tensile testor (TENSILON of Toyo Baldwin Co., Ltd.) at a chuck interval of 100 mm and a pulling rate of 10 mm/min in a room adjusted to a temperature of 20 DEG C and a relative humidity of 50 %. The Young's modulus was calculated from the tangent of a rising portion of the obtained stressdistortion curve.

(b) flexural rigidity

[0124] This was calculated based on the following equation from the inclination of the tangent of a rising portion of the obtained sites-distribing course by pulling the film at a chuck interval of 100 mm and a pulling rate of 5 mm/min using the same apparatus and measurement conditions as in the measurement of Young's modulus. Recural rigidity (mg mm) = {(10<5> (mg/kg) x (inclination of tangent (kg/mm<2>)) x (film thickness /mm)>3/12

(c) tensile strength at break

[0125] A stress at the time when the film was broken by pulling at a chuck interval of 100 mm and a pulling rate of 100 mm/min using the same apparatus and measurement conditions as in the measurement of Young's modulus was taken as break strength.

(8) thermal shrinkage

[0128] Ten 350 mm x 350 mm samples of the film were prepared, but opinits were marked on a center option of the sample be ensure that a louge length (10 (mm)) between the two opinits should be 300 mm. Ten marked films were hung in a hot-air circulating constant temperature chamber maintained at 150 DEG Courder no transion, these films were taken out from the chamber later they were held in the chamber for 30 minutes, and the length (1, (mm)) between the two points of the films was measured. Thermal shimkage thermal shinkage a 150 x (160 - 1100).

(9) winding property

[0127] After winding conditions at the time of slitting were optimized, the film was slit to obtain 10 rolls, each having a width of 550 mm and a length of 9,000 m and the rolls were then left for 1 week. The whichgy properly of the rolls was evaluated based on the following criteria from the number of rolls which could be commercialized judging from occurrence of film winkles.

<tb>

<tb>Head Col 1: number of commercializable rolls

<tb>Head Col 2 criteria

<tb>9 or more<SEP>O &cir& <tb>7 to 8<SEP> &cir&

<tb>4 to 6<SEP>x

<tb>3 or less<SEP>xx

<tb></TABLE>

(10) abrasion resistance (calendering)

[0128] This was evaluated using a three-roll mini-super calender (nylon roll x steel roll). The film was caused to run 200 m at a speed of 50 m/min at processing temperature of 80 DEG C and a linear pressure of 200 kg/cm. This was judged from abrasion dust adhered to the nylon roll of the calender based on the following criteria

<tb><TABLE> Columns=2

```
<tb>no abrasion dust on nylon roll<SEP>O &cir&
<tb>slight amount of abrasion dust on nylon roll<SEP> &cir&
<tb>large amount of abrasion dust on nylon roll<SEP>x
<tb></tb><tb></tb><tb></tb><tb></tb></tb></tb>
(11) electromagnetic conversion characteristics
[0129] A magnetic coating prepared as follows was applied to the surface of the film to a thickness of 1.2
mu m, aligned in a DC magnetic field of 2,500 Gauss, dried by heating at 120 DEG C and subjected to
super-calendering (linear pressure: 300 kg/cm, temperature; 90 DEG C), and the resulting film was wound
up into a roll. This roll was left in an oven at 55 DEG C for 3 days
aration of magnetic coating>
[0130] After the composition shown below was charged into a ball mill, kneaded for 16 hours and dispersed,
5 parts by weight of an isocyanate compound (Desmodule L of Bayer AG) was added to the ball mill and
dispersed by high-speed shearing for 1 hour to prepare a magnetic coating.
* composition of coating:
<tb><TABLE> Columns=2
<tb>needle-like Fe particles<SEP>100 parts
<tb>vinyl chloride-vinyl acetate copolymer (Slec 7A of Sekisul Chemical Co., Ltd.)<SEP>15 parts
<tb>thermoplastic polyurethane resin<SEP>5 parts
<tb>chromium oxide<SEP>5 parts
<tb>carbon black<SEP>5 parts
<tb>lecithin<SEP>2 parts
<tb>fatty acid ester<SEP>1 part
<tb>toluene<SEP>30 parts
<tb>methyl ethyl ketone<SEP>50 parts
<tb>cyclohexanone<SEP>70 parts
<tb></tb></tb>
[0132] Further, a coating solution having the following composition was applied to the film surface opposite
to the magnetic recording layer of the film to a thickness of 0.8 mu m as a back coal layer, dried and cut to
give a magnetic tape
* composition of back coat laver
<tb><TABLE> Columns=2
<tb>carbon black<SEP>100 parts
<tb>thermoplastic polyurethane resin<SEP>60 parts
<tb>isocyanate compound<SEP>18 parts
<tb>(colonate L of Nippon Polyurethane Kogyo<SEP>Co., Ltd.)
<tb>silicone oil<SEP>0.5 part
<tb>methyl ethyl ketone<SEP>250 parts
<tb>toluene<SEP>50 parts
<tb></TABLE>
[0134] The characteristic properties of the tape were measured using the following commercially available
devices
* used devices:
[0135] 8 mm video tape recorder: EDV-6000 of Sony Corp.
[0136] S/N measurement: noise meter of Shibasoku Co., Ltd.
(a) S/N measurement
[0137] The S/N ratio of a magnetic video tape was measured and evaluated based on the following criteria
using a value in Comparative Example 3 as a reference
<tb><TABLE> Columns=2
<tb>larger than (reference value + 3 dB)<SEP>O &cir&
<tb>(reference value + 3 dB) &tilde& (reference value + 1 dB)<SEP> &cir&
<tb>smaller than (reference value + 1 dB)<SEP>x
<tb></TABLE>
(b) running durability
[0138] The tape was caused to run under high temperature and high humidity conditions of a temperature of
55 DEG C and a humidity of 75 %RH for continuous reproduction, a reproduced image was observed, and
running durability was judged from the vibration of the image.
<tb><TABLE> Columns=2
<tb>image is not vibrated<SEP> &cir&
<tb>image is vibrated<SEP>x
```

<tb></TABLE>

#### Example 1

[0139] 25 md/s of parapheny/enddamme and 25 md/s of 24-diaminocipiency letter as a mime components and 50 md/s of terephishica acid obtinide as an acid components were polymerized in NMP (N-methylgroridione). Meanwhile, calcium hydroxide having an average particle diameter of 8,000 mm (a product of noise Sekkal Co. L. Li) was desperted in NMP 9, be indocepitately, round to an average particle diameter of 2,300 mm (by a bindepender) of 50 mm on meshes manufactured by Whom Pail L. Li) to peptie are NMP 34 may of caction in plottode having an average particle diameter of 2,300 mm (by a solid production of 50 mm) of 10 md/s of terephishilat cacid tolknick wheneys a stock solution for film of 10 md/s of terephishilat cacid tolknick wheneys a stock solution for film of 10 md/s of terephishilat cacid tolknick wheneys a stock solution for film of 10 md/s of 10 md/s of terephishilat cacid tolknick wheneys a stock solution for film of 10 md/s of 10 md/s of terephishilat cacid tolknick wheneys a stock solution for film of 10 md/s of 10 md/s

[0:44] The obtained stock solution for film formation was heated at 100 DEG C, cast onto a metal bett heated at 100 DEG C form a nazzale heated at 100 DEG C for almost send network at 100 DEG C for almost send heated stoppus at 120 DEG C and then 150 DEG C to be dired for a total of 10 minutes so as to give an uncreasted film having self-holding propriets. This uncreinted film was separated from the best continuously and introduced into a water tank to remove the solvent and the salt, and then dried at 150 DEG C for 15 minutes.

[0141] The obtained unoriented film was stretched to 2.5 times between low-speed and high-speed rolls at a 14th Timeprature of 35 DEG. can dit her supplied to a stetler to be stretched to 3 times at 400 DEG. to give a blaxalily oriented film. This blaxially oriented film was heet set at 400 DEG. C for 1 minute to give an aromatic powlamide film having a final this/cines of 4.0 m un.

[0142] The average particle diameter of the residual neutralizing agent contained in the obtained film was 300 nm. Other physical properties and characteristic properties of the film are shown in Table 1.

#### Evample 2

[0143] Calcium hydroxide which was the same neutralizing agent as in Example 1 was ground to an average particle dameter of 4,800 m by a sand grinder and then, on NMP slurry was prepared directly. This slurry was added to the above polymer to ensure that an amount of calcium hydroxide became 50,06 molf/s based on 50 molf/s of teraphthalic acid chloride, to prepare a stock solution A for film formation (p.H. 52. Loadnthmic vascolary of the observed 3.5.

[D144] Maximp January Britan (1944) Mary Britan

[0145] Further, these stock solutions A and B were mixed together in equal amounts to prepare a final stock solution, and an aromatic polyamide film was obtained in the same manner as in Example 1.

[0146] The particle size distribution curve of the residual neutralizing agent contained in the obtained film had peaks at 500 nm and 180 nm. The physical properties and characteristic properties of the film are shown in Table 1

### Example 3

(10.4T) Calcium hydroxide which was the same neutralizing agent as in Example 1 was ground to an average particle immeter of 2,000 m by a sand prinder and then, an NMP silvin was prepared directly. An eromatic polyemide film was obtained in the same menore as in Example 1 except that this surry was added to the above opymer to ensure that an amount of calcium hydroxide beame \$2.3 mol% blased on 50 mol% of breighthalic acid chioride, to prepare a stock solution for film formation (pil-14.8, logarithms vaccounty of the polymer 3.5). The physical properties and characteristic properties of the film are shown in

## Example 4

[0.145] Calcium hydroxide wishch was the same neutralizing agent as in Example 1 was ground to an wertage particle interest of 2.00 on they a sand grinder and filtered with a filter (HDCII of 2.0 ms in meables remarked used by Almach Pall Ltd.) to prepare a media Barry of disclaim hydroxide having an everage profice along the properties of the properties and characteristic properties of the filter graphithm by dozenty of the polymer 2.0 The physical properties and characteristic properties of the filter properties of the properties of 2.0 The physical properties and characteristic properties of the filter properties of the properties of 2.0 The physical properties and characteristic properties of the filter properties of the properties of the properties of the properties and characteristic properties of the filter properties of the properties of the properties of the properties and characteristic properties of the filter properties of the prop

### Example 5

(0.149) Calcium hydroxide which was the same neutralizing agent as in Example 1 was ground to an average particle diameter of 2.000 mm by a sand grinder and filtered with a filter (HDCI of 2.00 mm by an manufactured by Nihon Pall Ltd.) to prepare an NMP sarry of calcium hydroxide having an average particle of 600 mm. An aromatic polyamide film was obtained in the same manner as in Example 1 except that this slurry was added to the above polymer to ensure that an amount of calcium hydroxide became 57 3 mol% based on 50 mol% of terephthalic acid chloride, to prepare a stock solution for film formation (pH: 5.0. logarithmic viscosity of the polymer, 3.5). The physical properties and characteristic properties of the film are shown in Table 1

## Example 6

[0150] An aromatic polyamide film was obtained in the same manner as in Example 1 except that the film was stretched to 2.0 times in a longitudinal direction and to 3.5 times in a transverse direction. The physical properties and characteristic properties of the film are shown in Table 1.

## Example 7

[0151] An aromatic polyamide film was obtained in the same manner as in Example 1 except that the film was stretched to 2.8 times in a longitudinal direction and to 2.8 times in a transverse direction. The physical properties and characteristic properties of the film are shown in Table 1.

#### Example 8

[0152] An aromatic polyamida film was obtained in the same manner as in Example 1 except that calcium carbonate was used as a neutralizing agent. The physical properties and characteristic properties of the film are shown in Table 1.

## Example 9

[0153] An aromatic polyamide film was obtained in the same manner as in Example 1 except that magnesium hydroxide was used as a neutralizing agent. The physical properties and characteristic properties of the film are shown in Table 1

#### Example 10

[0154] An aromatic polyamide film was obtained in the same manner as in Example 1 except that the molar ratio of paraphenylenediamine to 3,4'-diaminodiphenyl ether as amine components was changed to 37.5/12.5 (total of 50 mol%). The physical properties and characteristic properties of the film are shown in Table 1

# Example 11

[0155] An aromatic polyamide film was obtained in the same manner as in Example 1 except that the molar ratio of paraphenylenediamine to 3.4'-diaminodiphenyl ether as amine components was changed to 12.5/37.5 (total of 50 mol%). The physical properties and characteristic properties of the film are shown in

## Comparative Example 1

[0156] An aromatic polyamide film was obtained in the same manner as in Example 1 except that a dried unoriented film was not stretched. The physical properties and characteristic properties of the film are shown in Table 2. The unoriented film had few protrusions on the surface, was inferior in slipperiness and winding property, and had low Young's moduli

## Comparative Evample 2

[0157] An aromatic polyamide film was obtained in the same manner as in Example 1 except that an NMP sturry of calcium hydroxide was added to a polymerization system to adjust pH to 3.8 so as to prepare a stock solution for film formation. Particles regarded as the residual neutralizing agent were not found in the obtained film. Other physical properties and characteristic properties of the film are shown in Table 2.

## Comparative Example 3

[0158] An aromatic polyamide film was obtained in the same manner as in Example 1 except that a stock solution for film formation (pH 2.3) was prepared without adding an NMP slurry of a neutralizing agent to a polymerization system. Particles regarded as the residual neutralizing agent were not found in the obtained film. Other physical properties and characteristic properties of the film are shown in Table 2

### Comparative Example 4

[0159] An aromatic polyamide film was obtained in the same manner as in Example 1 except that an NMP sturry of calcium hydroxide was added to a polymerization system to adjust pH to 3.8 and a pre-dispersed

silica having an average particle diameter of 10 nm was further added to prepare a stock solution for film formation. Particles regarded as the residual neutralizing agent were not found in the obtained film. Other physical properties and characteristic properties of the film are shown in Table 2.

#### Comparative Example 5

[0160] An aromatic polyaminis firm was obtained in the same manner as in Example 1 except that an NIMP stury of calculm inferrode was adord to a polymerization by system to adjust pit 1.63 and a pre-religional silica having an average particle diameter of 4.30 mm was further adord to prepare a stock solution for film formation. Particles regarded as the racidiant neutralizing agent were not found in the obtained film. Other physical properties and characteristic properties of the film are shown in Table 2.

EMI42.1

FMI43.1

EMI44.1

#### Example 12

[1011] Zs moll's of paragheny/enediamine and 25 moll's of 3.4 -diamnosighenyl ether as anima components and 50 molly of terephishic and dictionide as an audic component were polymerized in NNP (k-methylpyrrolidone). Meanwhile, calcium hydroxide having an average partical diameter of 8.000 nm (control of 10.000 nm) and the product of incurse Sekstali Co. Ltd. Vasa dispersari of NMP by a homogenzer; product of an average partical diameter of 1.000 nm by a send ginder and filteral with a filter (FIDCI of 5 min meether annualscutered of 200 nm) and the prepara on NMP author of calcium hydroxide having an average partical diameter of 200 nm by a send ginder and filteral with a filter (FIDCI of 5 min meether annualscutered of 200 nm). The prepara on NMP author of calcium hydroxide having an average partical diameter of 200 nm) and the preparation of the component of the control of the component of the control of t

[0162] The obtained stock solution was heated at 100 DEG C, cast onto a metal belt heated at 100 DEG C from a nozzle heated at 100 DEG C, oried at 100 DEG C for 2 minutes and heated stepwise at 120 DEG C and then to 150 DEG C to be dried for a total of 10 minutes so as to give an unoriented film having self-holding properties. This unoriented film was separated from the belt continuously and introduced into a water tank to remove the solvent and the salt and the noride at 150 DEG C for 15 minutes.

[0163] The obtained unoinented film was stretched to 2.5 times between low-speed and high-speed rolls at a film temperature of 350 DEG C, and then supplied to a stenter to be stretched to 3.0 times at 400 DEG C to give a biaxially oriented film. This biaxially oriented film was heat sat at 400 DEG C for 1 minute to give an aromatic polymide film having a final thickness of 3.0 m cm.

[0164] The average particle diameter of the residual neutralizing agent contained in the obtained film was 60 nm. Other physical properties and characteristic properties of the film are shown in Table 3.

## Example 13

[9165] Calcium hydroxides which was the same neutralizing agent as in Example 12 was ground to an average particle deniented 2.000 nm by a sand grinder and filtered with a filter (HOLI of 10 nm um meables manufactured by Rilton Pall Ltd.) to prepare an NRP allry of Goldam yddoxide having an average particle that this satury was added to the above opywher to ansure that a manuroff calcium hydroxide bearing 50.15 m/dr. Spatial particles and the particles of t

# Example 14

[0166] Calcium hydroxide which was the same neutralizing agent as in Exemple 12 was ground to an average particle desired 2.00 of not her and grant produced in the Children of white filter (Ph.Children of the many of the Children of the same produced in the control of the control of the children of th

### Example 15

[0167] Calcium hydroxido which was the same neutralizing agent as in Example 12 was ground to an average particle dismeter of 2,000 m hy a sand printer and filtered with a filter HOLT of 20 m. m neutres manufactured by Nhon Pall Ltd.) once by pepare an NMP altrny of calcium hydroxide having an average particle diameter of 300 m. n. an aromatic polyratine film was obtained in the same manner as in Example 12 except that this sturry was added to the above polymer to ensure that an amount of calcium hydroxide 12 except that this sturry was added to the above polymer to ensure that an amount of calcium hydroxide became \$0.1 molth based on 50 molths' of receptibal case of all-foliorised to pepare as sock solution for film formation (p.H. S.2. logar/miner viscosity of the polymer 3.5). The physical properties and characteristic properties of the film are shown in Table .

#### Example 16

[0168] Calcium hydroxide which was the same neutralizing agent as in Example 12 was ground to an average particle diameter of 2,000 nm by a said ginder and filtered with a filter (HDCI of 15 nm meshes manufactured by Nihon Pall Ltd.) to prepare an NMP sturry of calcium hydroxide having an average particle diameter of 550 nm. An aromatic polyamide film was obtained in the same manner as in Example 12 except that this slurry was added to the above polymer to ensure that an amount of calcium hydroxide became 50.6 mol% based on 50 mol% of terephthalic acid dichloride to prepare a stock solution for film formation (pH: 5.3, logarithmic viscosity of the polymer: 3.5). The physical properties and characteristic properties of the film are shown in Table 3.

### Example 17

[0169] An aromatic polyamide film was obtained in the same manner as in Example 12 except that the film was stretched to 2.0 times in a longitudinal direction and to 3.5 times in a transverse direction. The physical properties and characteristic properties of the film are shown in Table 3

#### Example 18

[0170] An aromatic polyamide film was obtained in the same manner as in Example 12 except that the film was stretched to 2.8 times in a longitudinal direction and to 2.8 times in a transverse direction. The physical properties and characteristic properties of the film are shown in Table 3.

## Example 19

[0171] An aromatic polyamide film was obtained in the same manner as in Example 12 except that calcium carbonate was used as a neutralizing agent. The physical properties and characteristic properties of the film are shown in Table 3.

#### Evample 20

[0172] An aromatic polyamide film was obtained in the same manner as in Example 12 except that magnesium hydroxide was used as a neutralizing agent. The physical properties and characteristic properties of the film are shown in Table 3.

## Example 21

[0173] An aromatic polyamide film was obtained in the same manner as in Example 12 except that the molar ratio of paraphenylenediamine to 3,4'-diaminodiphenyl ether as amine components was changed to 37.5/12.5 (total of 50 mol%). The physical properties and characteristic properties of the film are shown in Table 3

# Example 22

[0174] An aromatic polyamide film was obtained in the same manner as in Example 12 except that the molar ratio of paraphenylenediamine to 3,4"-diaminodiphenyl ether as amine components was changed to 12 5/37.5 (total of 50 mol%). The physical properties and characteristic properties of the film are shown in Table 3

# Comparative Example 6

[0175] An aromatic polyamide film was obtained in the same manner as in Example 12 except that a dried unoriented film was not stretched. The physical properties and characteristic properties of the film are shown in Table 4. This uncriented film had few protrusions on the surface, was inferior in slipperiness and winding property and had low Young's moduli. Therefore, the film was inferior in electromagnetic conversion. characteristics

## Comparative Example 7

[0176] An aromatic polyamide film was obtained in the same manner as in Example 12 except that an NMP slurry of calcium hydroxide was added to a polymerization system to adjust pH to 3 8 so as to prepare a stock solution for film formation. Particles regarded as the residual neutralizing agent were not found in the obtained film. Other physical properties and characteristic properties of the film are shown in Table 4

### Comparative Example 8

[0177] An aromatic polyamide film was obtained in the same manner as in Example 12 except that a stock

solution for film formation (pH was 2.3) was prepared without adding an NMP sturry of a neutralizing agent to a polymerization system. Particles regarded as the residual neutralizing agent were not found in the obtained film. Other physical properties and characteristic properties of the film are shown in Table 4.

## Comparative Example 9

[0178] An aromatic polyamide film was obtained in the same manner as in Example 12 except that an unoriented film separated from a netal belt was died at 200 DEG C for 30 minutes without desalting in a water tank. The physical properties and characteristic properties of the film are shown in Table 4.

## Comparative Example 10

(0.179) An aromatic polyamide film was obtained in the same manner as in Example 12 except that an NIMP sharing of calcum syntrotice was added to a polymerization system to adjust pit 0.3 and any peri-deperated silica having an average particle diameter of 10 nm was further added to prepare a stock solution for film formation. Particles registed as the residual neutralizing agent were not found in the obtained film. Other physical properties and characteristic properties of the film are shown in Table 4.

```
EMI52.1
EMI53.1
```

EMI54.1

[0180] In above Table 3 and Table 4, halogen detection, friction coefficient, pin abrasion, C/N, running durability and moisture heat stability were measured and defined by the following methods.

## (12) detection of halogen ions on film surface

[0181] A 10 mm x 10 mm sample of the film was measured using an X-ray photoelectric spectrophobmeter (ESCALAR-200 of VG Co., Ltd.) and an Mg-4 sighal ine as an excitation X ray. The detection of halogen loss (Fe->, CI<->, Br<->, I<->) was judged from the obtained spectrum based on the following criteria -Lb>-₹7ABLE-5 Columns=2

```
<tb>no halogen ions are detected<SEP> &cir&
<tb>halogen ions are detected<SEP>x
```

<tb></TABLE>

[0182] As for the detection of halogen ions, CI<> ions are considered to be existent when a peak derived from a 2p-orbit appears at a binding energy of less than 203 eV. As a peak derived from covalent CI atoms which are not free CI<> ions appears at a round 203 to 25 eV, it is distinguished from the above peak.

## (13) friction coefficient

<tb></TABLE>

(0183) A static friction coefficient (mus) under a load of 1 kg was measured in accordance with ASTM D1894-63 using a slippory measuring instrument (manufactured by Toyo Testoc, Ltd.) and a glass plate <a href="https://doi.org/10.1016/j.chm/s-7.4016-0.0016/j.chm/s-7.4016-0.0016/j.chm/s-7.4016-0.0016/j.chm/s-7.4016-0.0016/j.chm/s-7.4016-0.0016/j.chm/s-7.4016-0.001

# (14) abrasion resistance (against pin)

[0184] Abrasion resistance was measured using an apparatus shown in Fig. 1 as follows. In Fig. 1, reference numeral 1 denotes a feed red. 2 a lension controller, 3, 5, 6, 8, 9 and 11 free rollers, 4 a tension detector (nilet), 7 stainless steel SUS304 fixing pin (outer diameter 5 mm, surface roughness Ra 20 nm), 10 a tension detector (outlet), 12 a guide roller and 13 a take-up redi.

[0185] At a temperature of 20 DEG C and a humidity of 80 %RH, the film cut to a width of 1/2 inch was brought into contact with the fixing pin 7 at angle of 8 thetas, = (80/180) pi radian (80 DEG) and caused to run 100 m at a speed of 2 m/min (relat tension was fixed at 40 g), and abrasion resistance was judged based on the following criteria.

```
<tb>No abrasion dust is adhered to the pin<SEP>O &cir&

<tb>SEP> &cir&

<tb>Acir&

<tb>Acir</tb>
<tb>Acir
```

-to- 417 to ac-

# (15) electromagnetic conversion characteristics

(2)88] Two 100 % cobalt ferromagnets thin film layers were formed on the surface of the film to have a total trickness of 90 g. zum deach typer that a thickness of 400 g. zum deach typer that a thickness of 400 g. value (10 tim grey and a fluorene-containing cathorytic acid-based butricant layer were formed sequentiating on the above surface of the thin film layers and a fluorene duty the formed sequentiating on the above surface of the thin film layers and a fluor cost layer was formed on the surface opposite to the magnetic layer by a horse method. Thereafter, the resulting film was all tim a 8 mm which was then loaded in an a commercially available 8 mm which casasetts. Then be characteristic properties of this tape were measured using the following commercially available devices used devices.

8 mm video tape recorder EDV-6000 of Sony Corp.

C/N measurement: noise meter of Shibasoku Co., Ltd.

#### (a) C/N measurement

[0.187] A signal having a recording wavelength of 0.5 mm (frequency of about 7.4 MHz) was recorded; the ratio of values of its reproduction signal at 6.4 MHz and 7.4 MHz, was taken as CIV.0 of the tape, the CIV.0 of deposited tape for a commercially available 8 mm video was regarded as 0.4B, and CIV. was evaluated with a relative value based on the following criteria.

<tb><TABLE> Columns=2 <tb>larger than (reference + 5 dB)<SEP>O &cir&

<tb>(reference + 5 dB) to (reference + 1 dB)<SEP> &cir&

<tb>smaller than (reference + 1 dB)<SEP>x</tb>

<tb></TABLE>

## (b) running durability

[0188] The tape was caused to run at a temperature of 55 DEG C and a humidity of 75 %RH for continuous reproduction, a reproduced image was observed, and running durability was judged according to the wibration of the image

<tb><TABLE> Columns=2

<tb>!mage is not vibrated<SEP> &cir& <tb>!mage is vibrated<SEP>x

<tb>Image is vibrated<SEP>x <tb></TABLE>

### (16) stability under moist/heat conditions

[0.188] After the tape was left to stand under the conditions of a temperature of 60 DEG C and a humidity of 80 %RH for 60 hours, stability under moist and heat conditions was evaluated according to the corrosion of the deposited surface by visual inspection and the count value of defects produced by the corrosion of the deposited surface by observation through an optical microscope based on the following criteria, the TABLE D-COMISMS OCCUMISMS.

<tb>

<tb>Head Col 1: visual inspection <tb>Head Col 2: microscope

stb>Head Col 3: criteria

<tb>no corrosion<SEP>less than 10/84 mm<2><SEP>O &cir&

<tb>no corrosion<SEP>10/64 mm<2> or more<SEP> &cir& <tb>corroder(<SEP><SEP>x

### Example 23

[0190] The unoriented film obtained in Example 1 was stretched to 2.5 times between low-speed and highposed rolls at a film memperature of 500 DEC and then supplied to a stentor to be stretched to 3.0 times at 400 DEC 1 to give a biasidity oriented film. This biaxisity oriented film was heat set at 400 DEC 0 for 1 minute, cocede to room temperature, coacied with the following coasing solution [ii] in a mount of 1.0 g/m22 using a roll coater (coating film thickness after drying; 10 mm), and dired with hot air at 210 DEC for 30 seconds to you an easily advise bibliosally oriented durinate polyamide film having a final film.

[0191] The average particle diameter of the residual neutralizing agent contained in the obtained film was 300 nm. Other physical properties and characteristic properties of the film are shown in Table 7.

(0192) The coating solution [1] was prepared as follows:

cproduction of copolyester resin A>

(1913) 30 parts by weight of dimethyl 2.5-maphthalenedicarboxylic acid. 6 parts by weight of dimethyl sophimalist, 4 parts by weight of sold minetyl sophimalists, 4 parts by weight of sold parts by weight of sold parts by weight of sold parts by weight of an addition product of baphenic A with ethylene outsid (represented by the following sold parts by weight of an addition product of bapt parts of a weight of the sold productions of the sold parts of the so

(m + n = 4 (average value))

[0194] To this reaction system was then added 0.6 part by weight of Irganox 1010 (Ciba Geigy Co., Ltd.), temperature was deveated gradually to 255 DEG C, and pressure inside the system was reduced to 1 mmH<sub>2</sub> to carry out a polycondensation reaction to give a copolyester resin A having an intrinsic viscosity

(ocp) of 0.64 [0195] The composition of this copolyester resin A is shown in Table 5 preparation of polyester dispersion in water "a"> [0196] 20 Parts by weight of the copolyester resin A was dissolved in 80 parts by weight of tetrahydrofuran and 180 parts by weight of water was added dropwise to the obtained solution under high-speed agitation of 10,000 rpm to give a semi-opaque dispersion tinted with blue. This dispersion was distilled at a reduced pressure of 20 mmHg to remove tetrahydrofuran. Thus, a polyester dispersion in water "a" having a solid content of 10 wt% was obtained preparation of coating solution fill for coating layer> [0197] A water dispersion composition comprising 10 parts by weight of a 10 wt% aqueous solution of polyoxyethylene nonylphenyl ether as a surfactant and 90 parts by weight of the obtained polyester dispersion in water "a" was diluted with water to prepare an aqueous coating solution for a coating layer (solid content: 1.0 wt%). The composition is shown in Table 6 [0198] Coating solutions [II] to [VI] used in the following examples and comparative examples were prepared as follows preparation of coating solutions [II] to [VI] for coating layer> [0199] Copolyesters B to F were prepared in the same manner as described above except that the compositions of the copolymers shown in Table 5 were used to prepare the corresponding water dispersions "b" to "f" in the same manner as described above. Coating solutions [ii] to [Vi] for coating layers were prepared using these water dispersions according to formulations shown in Table 6.

<tb><TABLE> Id=Table 5 Columns=7 <tb>composition of copolvester for coating laver(mol%) <tb>Head Col 1: polyester <tb>Head Col 2 to 4 AL=L. dicarboxylic acid components <tb>Head Col 5 to 6 AL=L: glycol components <tb>Head Col 7 AL=L: dispersion in water (10%) <tb>SubHead Col 1: <tb>SubHead Col 2: NDCA <tb>SubHead Col 3: IPA <tb>SubHead Col 4: NSIPA <tb>SubHead Col 5 EG <tb>SubHead Col 6: BPA-P <tb><SEP>A<SEP>90<SEP>6<SEP>4<SEP>70<SEP>40<SEP>a <tb><SEP>B<SEP>98<SEP>-<SEP>2<SEP>90<SEP>10<SEP>b <SEP>6<SEP>60<SEP>38<SEP>2<SEP>100<SEP>-<SEP>c <tb><SEP>D<SEP>60<SEP>32<SEP>8<SEP>70<SEP>30<SEP>d <tb><SEP>E<SEP>30<SEP>68<SEP>2<SEP>100<SEP>-<SEP>e <tb><SEP>E<SEP>60<SEP>38<SEP>2<SEP>30<SEP>70<SEP>4 NDCA: 2 6-paphthalanadicarbovylic acid IPA: isophthalic acid NSIPA: 5-sodium sulforsonhthalic acid EG: ethylene glycol BPA-P: addition product of bisphenol A with propylene oxide <tb></TABLE> <tb><TABLE> Id=Table 6 Columns=5 <tb>composition of coating solution for coating layer <tb>Head Col 1: <tb>Head Col 2 to 3: polyester <tb>Head Col 4: weight ratio of surfactant <tb>Head Col 5: concentration of coating solution (wt%) <tb> <tb>SubHead Col 1: <tb>SubHead Col 2: kind <tb>SubHead Col 3: weight ratio <br/>
<br <tb><SEP>IIII<SEP>b<SEP>90<SEP>10<SEP>1 <tb><SEP>(III)<SEP>c<SEP>90<SEP>10<SEP>1 <tb><SEP>(IV)<SEP>d<SEP>90<SEP>10<SEP>1 <tb><SEP>(VI<SEP>e<SEP>90<SEP>10<SEP>1 <br/>
<br Surfactant: polyoxyethylene nonylphenyl ether <tb></TABLE>

### Example 24

[0200] The procedure of Example 23 was repeated for the film of Example 2. The particle size distribution curve of the residual neutralizing agent contained in the obtained film had maximum peaks at 500 nm and

180 nm. Other physical properties and characteristic properties of the film are shown in Table 7.

#### Example 25

[0201] Calcium hydroxide winch vast the same neutralizing agent as in Example 23 was ground to an warrage particle disenser of 2,000 mm by a sand grinder and frilered with a filter (MPIOI of 20 mm m meshes manufactures by Ninon Pall Ltd.) to prepare an NNIP sturry of calcium hydroxide having an average particle disenser of 000 mm. As aromatic polyamide film was obtained in the same manner as in Example 23 except 30.4 mm) filter and the study of the same study of the same

[0202] The average particle diameter of the residual neutralizing agent contained in the obtained film was 180 nm. Other physical properties and characteristic properties of the film are shown in Table 7.

#### Evample 26

[0203] The procedure of Example 23 was repeated for the film of Example 3. The average particle diameter of the residual neutralizing agent contained in the obtained film was 600 nm. Other physical properties and characteristic properties of the film are shown in Table 7.

## Example 27

[0204] The procedure of Example 23 was repeated for the film of Example 5. The average particle diameter of the residual neutralizing agent contained in the obtained film was 300 nm. Other physical properties and characteristic properties of the film are shown in Table 7.

#### Evample 28

[0205] An aromatic polyamide film was obtained in the same manner as in Example 24 except that a dried unoriented film was stretched to 2.0 times in a longitudinal direction and to 3.5 times in a transverse direction. The physical properties and characteristic properties of the film are shown in Table 7.

#### Example 29

[0206] An aromatic polyamide film was obtained in the same manner as in Example 24 except that a dried unoriented film was stretched to 2.8 times in a longitudinal direction and to 2.8 times in a transverse direction. The physical properties and characteristic properties of the film are shown in Table 7.

## Examples 30 and 31

[0207] Aromatic polyamide films were obtained in the same manner as in Example 24 except that the coating solutions [I] and [III] were used in Examples 30 and 31, respectively. The physical properties and characteristic properties of the film are shown in Table 7.

### Comparative Example 11

[0208] An aromatic polyamide film was obtained in the same manner as in Example 23 except that a dried unonented film was not stetched. The physical properties and characteristic properties of the film are shown in Table 5. This unonented film had few protrusions on the surface, was lefterior in alligoriness and winding property and had low Young's moduli. Therefore, the film was inferior in electromagnetic conversion characteristics.

# Comparative Example 12

(2009) An aromatic polyamide film was obtained in the same manner as in Exemple 23 except that an NMP altury of calcium hydroxide was added to a polymerization system to adjust pirt to 3.8 as as to prepare a stock solution for film formation. Particles regarded as the reaction incubation; agent were not found in the obtained film. Other physical properties and characteristic properties of the film are shown in Table 8. This film having no protisions on the surface was inferior in blocking resistion.

### Comparative Example 13

(2010) An aromatic polyamide film was obtained in the same manner as in Example 23 except that as NME stury of calcium injectived having an example particle disenter of 13.00 nm which was prepared without being pricing by a sand grinder was added to a polymerization system to ensure that an amount of calcium hydrounde became 30 and/b based on 5 mmCnf of tree-brilland; acid dichorder. The average particle diameter of the residual requirationing agent contained in the obtained film was 2,250 mm. Other physical particles contained for the film were obtained the same of the interest invention, askagen registrating was particles contained in the film were obtained the rance of the interest invention, askagen registrating was unsatisfactory and appropriate surface smoothness was lost. Therefore, the film was inferior in electromagnetic conversion characteristics Comparative Examples 14 to 16 "[0211] Aromatic polyamide films were obtained in the same manner as in Example 24 except that coating solutions [IV] to [VI] for coating layers were used in Comparative Examples 14 to 16, respectively. The physical properties and characteristic properties of the films are shown in Table 8. EMISS 1 EMI66 1 EMI67.1 EMI68 1 [0212] in the above Examples 23 to 31 and Comparative Examples 11 to 16, the following physical property values were measured and defined by the following methods. (17) average particle diameter of colloidal particles added to adhesive layer [0213] The average particle diameter of colloidal particles was calculated using an image analyzer (Luzex 500 of Nippon Regulator Co., Ltd.) by photographing each particle at a magnification corresponding to the size of each particle with a scanning electron microscope. (18) coating thickness [0214] A small piece of the film was encapsulated in an epoxy resin (Epomount of Refinetec Co., Ltd.) and sliced to a thickness of 60 nm together with the resin using the Microtome 2050 of Reichert-Jung Co., Ltd. The slice was observed through a transmission electron microscope (H-800 of Hitachi, Ltd.) and the interface of the coating layer was searched for to obtain the thickness of the coating layer. (19) density of protrusions [0215] The above method (5) or the following method (19) was used according to a sample to be inspected Methods used in Examples and Comparative Examples are shown in Table 7. [0216] 25 Photos of the surface of the film were taken at random at a magnification of 50X using an optical microscope to count protrusions on the surface. The number of protrusions per 1 mm<2> was calculated from the average of measurement values. (20) surface roughness (WRa, WRz) [0217] The center plane average roughness (WRs) and the 10-point average roughness (WRz) were obtained using a surface analyzing software built in the non-contact 3-D roughness meter (TOPO-3D) of WYKO Co. Ltd. by conducting measurement under conditions of a 242 mu m x 239 mu m (0.058 mm<2>) measurement area at 10 or more sites and at a measurement magnification of 40X. a) center plane average roughness (WRa) [0218] This is a value calculated from the following expression and output FMI70.1 [0219] Zjk is a height on a 2-D roughness chart at a j-th position and a k-th position in a measurement direction (242 mu m) and a direction perpendicular to the direction (239 mu m) when these directions are divided into M and N sections, respectively. (b) 10-point average roughness (WRz) [0220] The average roughness WRz is obtained by averaging the roughnesses of the first to fifth highest

(21) abrasion resistance (calendering)

peaks (Hp) and the first to fifth deepest valleys (Hv).

[0221] This was evaluated using a three rolls mini-super calender (nylon roll x steel roll). The film was caused to run 2.000 m at a speed of 50 m/min at a processing temperature of 80 DEG C and a linear pressure of 200 kd/cm. Abrasion resistance was indeed from abrasion dust adhered to the nylon roll of the

## (22) adhesion of magnetic layer (MAG)

[0222] The following magnetic coating was applied to the firm (coated with a primer). Scotch tage No 900 of product of 3M Co. Lid) measuring 16 Am min width and 5 m in length was effect by a manual load rail manner that an air subble was not included herebelveen, the tape was refer by a manual load rail and the second of t

preparation of magnetic coating for evaluation;

[0223] 4.35 parts by weight of a 40 writ solution of infocellulose (falses containing 25 % of isonopsanol) (Sil 3/2 of loaked) Co. L. L. (a) loaced writemer (or coaling), 3/2, bart by weight of a polyseter resin (identification) and coaled magnetic and powers are solved to the containing of the solved and the coaled to the coaled to a power to the coaled to the coaled to a power to the coaled to the coaled to a power to the coaled to the coaled

[0224] To 50 parts by weight of this mother solution were added 48 parts by weight of an addition reaction product between trimethylol propane and follyinghen dislocyanatic (Colonate L of Nippon Polyurothane Kogyo Co., Ltd.) and 6.25 parts by weight of butyl acetate to finally obtain a magnetic coasting having a concentration of 42.7 tw% for evaluation

[0225] Evaluation was carried out from peel strength based on the following criteria

- <tb><TABLE> Columns=2 <tb>more than 40 g/cm<SEP> &cir&
- <tb>more than 40 g/cm<SEP> &ci <tb>30 to 40 g/cm<SEP> DELTA
- <tb>less than 30 g/cm<SEP> DELTA
- <tb></TABLE>

(23) blocking under moist/heat condition

[0236] Two films were superposed together in such a manner that a treated side of one film and an untreated side of the other film were in contact with each other, and applied with a pressure of 8 kg/cm<2> in an environment of 80 DEG C and 80 %RH for 17 hours. These films were peeled off from each other and blocking with most heat was evaluated from its peeling (6/5 on in width).

[0227] Evaluation was made from peeling based on the following criteria

- <tb><TABLE> Columns=2
- <tb>less than 10 g/5 cm<SEP>O
- <tb>10 to 15 g/5 cm<SEP> DELTA <tb>more than 15 g/5 cm or rupture<SEP>x
- CHARLES OF CHI OF PURPLES

Example 32

[0228] The unominated film obtained in the course of Example 1 was stretched to 2.5 times between lowspeed and high-pader oils sat a film experienzar of \$50 DECC and then supplied to a stretter to be stretched to 3.0 times at 40.0 DEC of to give a biavially oriented film. This biavaily oriented film was heat set at 400 DEC of the Timulus, clocided to room temperature, coarded with the following coarling soution, VIDI an amount of 1.0 g/m-2\* using a roll coaler, and diried with hot air at 210 DEC of the 30 seconds to give an adhesive biaviative oriented aromatic polygamine film haiving a final thickness of 40 m m.

[0229] The average particle diameter of the residual neutralizing agent contained in the obtained film was 300 nm. Other physical properties and characteristic properties of the film are shown in Table 10.

[0230] The coating solution [VII] was prepared as follows

[0231] A water dispersion composition composition to yet weight of the above polyester dispersion in water "a", 3.5 parts by weight of a 10 wt% water dispersion of cross-linked acryl particles having an average particle dismeter of 20 mm and 11.5 parts by weight of a 10 wt% aqueous solution of polyoxyethylene nonylphenyl either was diluted with water to prepare an aqueous coating solution [Viii] for a coating layer (solid content of 10 wt%).

[0232] Coating solutions [VIII] to [XIX] used in the following examples and comparative examples were prepared as follows.

coating solutions [VII] to [XIX] for coating layers>

```
(0233) The copolyesters B to F listed in the above Table 5 were used to prepare the corresponding
dispersions in water "b" to "f" in the same manner as described above, and coating solutions [VIII] to [XIX]
for coating layers were prepared in accordance with formulations shown in Table 9
<tb><TABLE> Id=Table 9 Columns=8
<tb>composition of coating solution for coating layer
cth>Head Col 1:
<tb>Head Col 2 to 3: polyester
<tb>Head Col 4 to 6, colloidal particle
<tb>Haad Col 7: weight ratio of surfactant
<tb>Head Col 8: concentration of coating solution(wt%)
>
<tb>SubHead Col 1:
<tb>SubHead Col 2: kind
<tb>SubHead Col 3: weight ratio
<tb>SubHead Col 4 kind
<tb>SubHead Col 5: particle size(d)
<tb>SubHead Col 6 weight ratio
<tb><SEP>IVIII<SEP>a<SEP>100<SEP>acryl<SEP>20nm<phi>><SEP>3 5<SEP>11.5<SEP>1
<tb><SEP>[VIII]<SEP>a<SEP>100<SEP>acryl<SEP>20nm<ph; ><SEP>3.5<SEP>11.5<SEP>1.5
<SEP>[IX]<SEP>a<SEP>100<SEP>acryl<SEP>40nm<phi>><SEP>7.1<SEP>10.6<SEP>1
<tb><SEP>IXI<SEP>a<SEP>100<SEP>core-shell<SEP>25nm<phi>><SEP>2.3<SEP>11.4<SEP>1
<tb><SEP>[XI]<SEP>8<SEP>100<SEP>silica<SEP>17nm<ph; ><SEP>47<SEP>116<SEP>16<SEP>17nm
<tb><SEP>[XII]<SEP>a<SEP>100<SEP>slica<SEP>25nm<phi>><SEP>4.7<SEP>11.6<SEP>1
<tb><SEP>[XIII]<SEP>a<SEP>100<SEP>silica<SEP>8nm<phi>><SEP>4.7<SEP>11.6<SEP>1
<b><SEP>[XIV]<SEP>a<SEP>100<SEP>silica<SEP>67nm<phi><SEP>14<SEP>12.8<SEP>1
<tb><SEP>[XV]<SEP>b<SEP>100<SEP>acryl<SEP>20nm<phi ><SEP>3.5<SEP>11.5<SEP>1
<tb><SEP>[XVI]<SEP>c<SEP>100<SEP>acryl<SEP>20nm< phi ><SEP>3 5<SEP>11.5<SEP>1
<tb><SEP>(XVII)<SEP>d<SEP>100<SEP>acryl<SEP>20nm<ph; ><SEP>3.5<SEP>11.5<SEP>1
<tb><SEP>[XVIII]<SEP>e<SEP>100<SEP>acryl<SEP>20nm<phi><SEP>3 5<SEP>11.5<SEP>1
<SEP>(XIX)<SEP>f<SEP>100<SEP>acryl<SEP>20nmohi ><SEP>3 5<SEP>11 5<SEP>1
```

## acryl: polymethyl methacrylate-divinylbenzene cross-linked product

core-shell type: core (total diameter/core diameter = 1.25): polystyrene-divinylbenzene cross-linked product, shell: polymethyl methacrylate silica: colloidal silica

surfactant: polyoxyethylene nonviphenyl ether

<tb></TABLE>

## Example 33

[0234] The procedure of Example 32 was repeated for the film of Example 2. The particle size distribution curve of the residual neutralizing agent contained in the obtained film had maximum peaks at 500 nm and 180 nm. Other physical properties and characteristic properties of the film are shown in Table 10.

### Example 34

[0235] The procedure of Example 32 was repeated for the film of Example 4. The average particle diameter of the residual nautralizing agent contained in the obtained film was 180 nm. Other physical properties and characteristic corporaties of the film are shown in Table 10.

## Example 35

[0238] The procedure of Example 32 was repeated for the film of Example 3. The average particle diameter of the residual neutralizing agent contained in the obtained film was 800 nm. Other physical properties and characteristic properties of the film are shown in Table 10.

## Example 36

[0237] The procedure of Example 32 was repeated for the film of Example 5. The average particle diameter of the residual neutralizing agent contained in the obtained film was 300 nm. Other physical properties and characteristic properties of the film are shown in Table 10.

## Example 37

[0238] An aromatic polyamide film was obtained in the same manner as in Example 33 except that a dried unoriented film was stretched to 2.0 times in a longitudinal direction and to 3.5 times in a transverse direction. The physical properties and characteristic properties of the film are shown in Table 10.

## Example 38

[0239] An aromatic polyamide film was obtained in the same manner as in Example 33 except that a direct unoriented film was stretched to 2.8 times in a longitudinal direction and to 2.8 times in a transverse

direction. The physical properties and characteristic properties of the film are shown in Table 10

#### Example 39

[0240] An aromatic polyamide film was obtained in the same manner as in Example 33 except that calcium carbonate was used as a neutralizing agent. The physical properties and characteristic properties of the film are shown in Table 10.

## Example 40

[0241] An aromatic polyamide film was obtained in the same manner as in Example 33 except that magnesium hydroxide was used as a neutralizing agent. The physical properties and characteristic properties of the film are shown in Table 10.

#### Example 41

(0.242). An arcmatic polyamide film was obtained in the same manner as in Example 33 except that the molar ratio of paraphenylenetamine to 3.4 disance(plenyle there as a mine components was changed to 37.5/12.5 (total of 50 mol%). The physical properties and characteristic properties of the film are shown in Table 10.

## Example 42

(9243). An aromatic polyamide film was obtained in the same manner as in Example 33 except that the molar ratio of paraphenylenedismine to 3.4-disminosiphenyl either as anine components was changed to 12,5/37.5 (total of 50 mol%). The physical properties and characteristic properties of the film are shown in Table 10.

#### Evamples 43 and 44

[0244] Aromatic polyamide films were obtained in the same manner as in Example 33 except that the coating solutions [XV] and [XVI] for coating layers were used in Examples 43 and 44, respectively. The physical properties and characteristic properties of the film are shown in Table 10.

# Example 45

[0245] An aromatic polyamide film was obtained in the same manner as in Example 33 except that the coating solution [VIII] for a coating layer was used in an amount of 2.0 g/m<2>. The physical properties and characteristic properties of the film are shown in Table 10.

## Examples 46 to 49

[0246] Aromatic polyamide films were obtained in the same manner as in Example 33 except that the coating solutions [IX] to [XII] for coating layers were used in Examples 46 to 49, respectively. The physical properties and characteristic properties of the film are shown in Table 10.

## Comparative Example 17

(0247) An aromatic polyamide film was obtained in the same manner as in Example 32 except that a dried unoriented film was not stetched. The physical properties and characteristic properties of the film are shown in Table 1.7. This uncertented film had few protrousnes derived from particles contained in the film on the surface, was inferior in slippemess and winding property and had low Young's moduli. Therefore, the film was inferior in electromagnetic conversion characteristics.

## Comparative Example 18

[0249] An aromatic polyamide firm was obtained in the same manner as in Example 32 except that an NIP stury of cardium infloration was added to a polymerization paystem to adjust pit of 38 as as to prepare a stock solution for film formation Particles regarded as the residual neutralizing agent were not found in the obtained film. Other physical properties and otheractivestic properties of the film are shown in Table 11. Till film having no portrusions derived from the particles contained in the film on the surface was inferior in sulpperintess and winding property.

### Comparative Example 19

(0249) An aromatic polyamide film was obtained in the same manner as in Example 32 except that an NMP slurry of a naturalizing agent was not added to a polymerization system to prepare a stock solution for film formation (pt. 2.3) Particles recarded as the residual neutralizing agent were not found in the obtained

film. Other physical properties and characteristic properties of the film are shown in Table 11. This film having no protrusions on the surface was inferior in slipperiness and winding property.

Comparative Evample 20

(0250) An anomatic polyamide firm was obtained in the same manner as in Example 32 except that an NME stury of calcium hydrocid was added to a polymerazion system to adquist pin 0.3 and a pre-dappend silica having an average particle diameter of 10 nm was further added to prepare a stock solution for film formation. Farticles regarded as the residual noturalizing agent were not found in the obtained film of the physical properties and characteristic properties of the film are shown in Table 11. Since the extensibly added tradrices had resulficient fairfully for the polymer and agglomentate, the obtained film vas inferior in deliver from the agglomerates of the particles, the film was inferior in electromagnetic conversion.

Comparative Example 21

(0251) An aromatic polyamide film was obtained in the same manner as in Exemple 32 except that an NMP surry of calcium hydroxide was added to a polymerization system to adjust pit 10-38 and a pre-displace silica having an average particle diameter of 430 am was further added to propare a stock solution for film formation. Particles regarded as the residual notatization goal reven not found in the obtained film. Other physical proparties and characteristic properties of the film are shown in Table 11. Since the externally added particles had our sufficient affirm? for the polymer and anglomerated, the doctained thin was infection and the properties of the particles, the film was infection in electromagnetic conventions.

Comparative Examples 22 to 24

[0252] Aromatic polyamide films were obtained in the same manner as in Example 33 except that the coating solutions [XVIII] and [XIX] were used in Comparative Examples 22 to 24, respectively. The physical properties and characteristic properties of these films are shown in Table 11.

Comparative Examples 25 to 26

[0253] Aromatic polyamide films were obtained in the same manner as in Example 33 except that the coating solutions [XIII] and EVIVI were used in Comparative Examples 25 and 26, respectively. The physical properties and characteristic properties of these films are shown in Table 11.

EMI82.1

EM183.1

EMI84 1

EMI85.1

EMISS 1

EMI87.1

[0254] in Examples 32 to 49 and Comparative Examples 17 to 26, the following physical properties were measured and defined by the following methods, provided that (17), (18), (19), (20) and (22) are the same as described above

(24) agglomeration of colloidal particles

[0255] 20 Photos of the coated surface of an adhesive aromatic polyamide film were taken at a magnification of 50,000X by a scanning electron microscope to count the number of agglomerates of 10 or more filler particles in the adhesive layer, and the number of agglomerates per 10 mu m<2> was calculated This was evaluated based on the following criteria.

<tb><TABLE> Columns=2 <tb>less than 10/10 mu m<2><SEP> &cir&

<tb>less than 10/10 mu m<2><SEP> &cir& <tb>10 to 30/10 mu m<2><SEP> DELTA

<tb>more than 30/10 mu m<2><SEP>X <tb></TABLE>

(25) abrasion resistance (calendering)

[0258] This was evaluated using a 5-stage min⊩super calender (steel roll x steel roll). The film was caused to run 2,000 m at a speed of 50 m/min at a processing temperature of 80 DEG C and a linear pressure of 200 kg/cm. Abrasion resistance was judged from the adhesion of abrasion dust of the top roll of the

calender based on the following criteria.

-tb>=TABLE> Columns=2

-tb>=n abrasion dust on top roll<SEP>O &cir&

-tb>=n abrasion dust on top roll<SEP> &cir&

-tb>=lange amount of abrasion dust on top roll<SEP>X

-tb>=IABLE>

(26) blocking under high-pressure, wet and heat conditions

[0257] Two films were superposed together in such a manner that a treated side of one film and an untreated side of the other film were in contact with each other, and applied with a pressure of 150 kg/cm-29 under an environment of 60 DEC can do 9kH for 65 hours. These films were peeted from each other and blocking under highpressure, wet and heat conditions was evaluated from its peeting (g/5 cm) in until high peeting for the peeting for the

[0288] Evaluation was made from peeling based on the following criteria.

-tb><TABLE Columns=2

-tb>-tes than 10 g/5 cm<SEP> &cirk

-tb>-10 to 16 g/5 cm<SEP> DELTA

-tb>-more than 15 g/5 cm -> rupture<SEP>X

-tb>-<TABLE.

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